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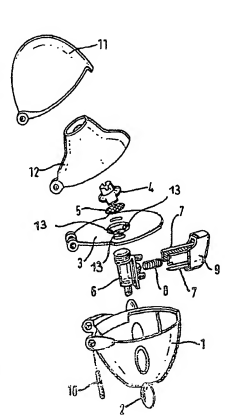
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[Continued on next page]

(54) Title: INHALATION KIT COMPRISUNG INHALABLE POWDER OF TIOTROPIUM

(57) Abstract: The invention relates to a method for the administration of powdered preparations containing tiotropium via inhalation.



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# INHALATION KIT COMPRISING INHALABLE POWDER OF TIOTROPIUM

The invention relates to a method for the administration of powdered preparations containing tiotropium by inhalation.

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# Background to the invention

Tiotropium bromide is known from European Patent Application EP 418 716 A1 and has the following chemical structure:

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Tiotropium bromide is a highly effective anticholinergic with a long-lasting activity which can be used to treat respiratory complaints, particularly COPD (chronic obstructive pulmonary disease) and asthma. The term tiotropium refers to the free ammonium cation.

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For treating the abovementioned complaints, it is useful to administer the active substance by inhalation. In addition to the administration of broncholytically active compounds in the form of metered aerosols and inhalable solutions, the use of inhalable powders containing active substance is of particular importance.

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With active substances which have a particularly high efficacy, only small amounts of the active substance are needed per single dose to achieve the desired therapeutic effect. In such cases, the active substance has to be diluted with suitable excipients in order to prepare the inhalable powder. Because of the large amount of excipient, the properties of the inhalable powder are critically influenced by the choice of excipient. When choosing the excipient its particle size is particularly important. As a rule, the finer the excipient, the poorer its flow properties. However, good flow properties are a prerequisite for highly accurate metering when packing and dividing up the individual doses of preparation, e.g. when producing capsules for powder inhalation or when the patient is metering the individual dose before using a multidose inhaler. It has also been found that the particle size of the excipient has a considerable influence on the proportion of active substance in the inhalable powder

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which is delivered for inhalation. The term inhalable proportion of active substance refers to the particles of the inhalable powder which are conveyed deep into the branches of the lungs when inhaled with a breath. The particle size required for this is between 1 and 10 µm, preferably less than 5 µm.

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Finally, it has been found that the intended therapeutic effect upon the administration of a pharmaceutical composition via inhaltion can be decisively influenced by the inhalation device.

Accordingly, the aim of the invention is to provide for a therapeutically efficient method for the administration of inhalable powders containing tiotropium. Another object of the invention is to provide for an inhalation kit comprising a tiotropium containing powder and an inhalation device, said kit being applicable in the method for administration mentioned before.

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### Detailed description of the invention

In the method according to the invention an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient is administered.

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Of particular interest for the method according to the invention is an inhalable powder containing 0.01 to 2 %, preferably 0.04 to 0.8 %, more preferably 0.08 to 0.64 % tiotropium in admixture with a physiologically acceptable excipient is administered. More preferably in the method according to the invention an inhalable powder containing 0.1 to 0.4 % tiotropium in admixture with a physiologically acceptable excipient is administered.

By tiotropium is meant the free ammonium cation. The counter-ion (anion) may be chloride, bromide, iodide, methanesulphonate, para-toluenesulphonate or methyl sulphate. Of these anions, the bromide is preferred.

Accordingly, the method according to the present invention preferably relates to inhalable powders which contain tiotropium in form of tiotropium bromide in an amount of 0.0012 to 6.02 %, in admixture with a physiologically acceptable excipient. Of particular interest for the method according to the invention is an inhalable powder containing 0.012 to 2.41 %, preferably 0.048 to 0.96 %, more preferably 0.096 to 0.77 % tiotropium bromide in admixture with a physiologically acceptable excipient is administered.

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More preferably in the method according to the invention an inhalable powder containing 0.12 to 0.48 % tiotropium bromide in admixture with a physiologically acceptable excipient is administered.

- Tiotroplum bromide is, depending on the choice of reaction conditions and solvents, obtainable in different crystalline modifications. Most preferred according to the invention are those powder preparations, that contain tiotropium in form of the crystalline tiotropium bromide monohydrate. Accordingly, the powdered preparations obtainable according to the invention preferably contain 0.0012 to 6.25 % crystalline tiotropium bromide monohydrate in admixture with a physiologically acceptable excipient is administered. Of particular interest for the method according to the invention is an inhalable powder containing 0.0125 to 2.5 %, preferably 0.05 to 1 %, more preferably 0.1 to 0.8 % crystalline tiotropium bromide monohydrate in admixture with a physiologically acceptable excipient is administered.
- More preferably in the method according to the invention an inhalable powder containing 0.12 to 0.5 % crystalline tiotropium bromide monohydrate in admixture with a physiologically acceptable excipient is administered.
- Examples of physiologically acceptable excipients which may be used to prepare the inhalable powders applicable according to the invention include, for example, monosaccharides (e.g. glucose or arabinose), disaccharides (e.g. lactose, saccharose, maltose), oligo- and polysaccharides (e.g. dextrane), polyalcohols (e.g. sorbitol, mannitol, xylitol), salts (e.g. sodium chloride, calcium carbonate) or mixtures of these excipients with one another. Preferably, mono- or disaccharides are used, while the use of lactose or glucose is preferred, particularly, but not exclusively, in the form of their hydrates, preferably in the form of their monohydrates.

In the method according to the invention the average particle size of the physiologically acceptable excipient is preferably between 10 to 500 µm, more preferably between 15 to 200 µm, most preferably between 20 to 100µm. If not otherwise emphazised the term average particle size according to the invention is to be understood as the Mass Median Aerodynamic Diameter (MMAD). Methods for the determination thereof are known in the art.

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Besides the coarser particle fraction of the excipient mentioned hereinbefore, the excipient can optionally additionally contain a specifically added fraction of excipient of finer particle size. This finer particle size fraction is characterized by an average particle size of 1 to 9 µm, preferably 2 to 8 µm, more preferably 3 to 7 µm. If a finer particle fraction is present the proportion of finer excipient in the total amount of excipient is 1 to 20 %, preferably 3 to 15%, more preferably 5 to 10%.

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When reference is made to a mixture within the scope of the present invention, this always means a mixture obtained by mixing together clearly defined components. Accordingly, when an excipient mixture of coarser and finer excipients is mentioned, this can only denote mixtures obtained by mixing a coarser excipient component with a finer excipient component.

The percentages given within the scope of the present invention are always percent by weight.

hereinbefore may effeciently be adminstered using inhalers that are characterized by a specific flow resistance (R).

The flow resistance of inhalers can be calculated via the following formula:

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$$V = \frac{1}{R} \cdot \sqrt{p}$$

wherein v is the volumetric flow rate (I/min),

p is the pressure drop (kPa), and

20 R is the flow resistance.

In the method according to the invention the flow resistance R characterising the inhaler is in a range of about  $0.01 - 0.1 \sqrt{kPa}$  min/l preferably in the range of about  $0.02 - 0.06 \sqrt{kPa}$  min/l.

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Accordingly, the invention relates to a method for the administration of an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m, and further characterized in that the said tiotropium containing powder is administered by an inhaler displaying a flow resistance of about 0.01 – 0.1  $\sqrt{kPa}$  min/l.

In another embodiment, the invention relates to a method for the treatment of airway diseases, particularly COPD (chronic obstructive pulmonary disease) and asthma, characterized in that an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m, is administered via inhalation by an inhaler displaying a flow resistance of about 0.01 – 0.1  $\sqrt{kPa}$  .min/l.

In another embodiment the invention relates to the use of an inhaler for the administration of a tiotropium containing inhalable powder via inhalation, characterised in that the inhalable powder contains tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m, and further characterized in that the said inhaler displays a flow resistance of about 0.01 – 0.1  $\sqrt{kPa}$  min/l.

In yet another embodiment the invention relates to an inhalation kit consisting of an inhaler displaying a flow-resistance of about  $0.01-0.1~\sqrt{kPa}$  min/l and an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm.

- In another preferred embodiment according to the invention the inhaler described in Figure 1 is applied. For the administration of tiotropium containing powders by inhalation by means of the inhaler according to figure 1, it is required to fill appropriate amounts of the powder into capsules. Methods for filling powders into capsules are known in the art.
- The inhaler according to figure 1 is characterised by a housing 1 containing two windows 2, a deck 3 in which there are air inlet ports and which is provided with a screen 5 secured via a screen housing 4, an inhalation chamber 6 connected to the deck 3 on which there is a push button 9 provided with two sharpened plns 7 and movable counter to a spring 8, a mouthpiece 12 which is connected to the housing 1, the deck 3 and a cover 11 via a spindle 10 to enable it to be flipped open or shut and three holes 13 with diameters below 1 mm in the central region around the capsule chamber 6 and underneath the screen housing 4 and screen 5.
  - The main air flow enters the inhaler between deck 3 and base 1 near to the hinge. The deck has in this range a reduced width, which forms the entrance slit for the air.
- Then the flow reverses and enters the capsule chamber 6 through the inlet tube. The flow is then further conducted through the filter and filter holder to the mouthpiece. A small portion of the flow enters the device between mouthplece and deck and flows then between filterholder and deck into the main stream. Due to production tolerances there is some uncertainty in this flow because of the actual width of the slit between filterholder and deck. In case of new or reworked tools the flow resistance of the inhaler may therefore be a little off the target value. To correct this deviation the deck has in the central region around the capsule chamber 6 and underneath the screen housing 4 and screen 5 three holes 13 with diameters below 1 mm. Through these holes 13 flows air from the base into the main air stream and

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reduces such slightly the flow resistance of the inhaler. The actual diameter of these holes 13 can be chosen by proper inserts in the tools so that the mean flow resistance can be made equal to the target value.

Accordingly, in a preferred embodiment the invention relates to a method for the administration of an inhalable powder containing tiotroplum, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm, by means of the inhaler according to figure 1, comprising

70 a housing, containing two-windows, a deck in which there are air inlet ports and which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring, a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in the central region around the capsule 15 chamber and underneath the screen housing and screen.

In another embodiment, the invention relates to a method for treatment of airway diseases, particularly COPD (chronic obstructive pulmonary disease) and asthma, charcterized in that an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm, is administered via inhalation by the inhaler according to figure 1, comprising

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a housing, containing two windows, a deck in which there are air inlet ports and 25 which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring, a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in the central region around the capsule chamber and underneath the screen housing and screen.

In another preferred embodiment the invention relates to the use of the inhaler according to figure 1, comprising a housing, containing two windows, a deck in which there are air inlet ports and which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring; a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in

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the central region around the capsule chamber and underneath the screen housing and screen,

for the administration of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm.

In yet another preferred embodiment the invention relates to an inhalation kit consisting of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm, and the inhalar according to figure 1, comprising

a housing, containing two windows, a deck in which there are air inlet ports and which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring, a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in the central region around the capsule chamber and underneath the screen housing and screen.

In another preferred embodiment according to the invention the inhaler according to 20 US 4,524,769 is applied. This inhaler (or inhalator) is activated by the air flow generated at inhalation. The disclosure of US 4,524,769 is incorporated herein by reference in its entirety.

Accordingly, in a preferred embodiment the invention relates to a method for the 25 administration of an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm, by means of the inhaler according to US 4,524,769, comprising a nozzle, a conduit connected to the nozzle, a storage chamber adjacent said conduit for storing said inhalable powder to be 30

dispensed by said inhalator, a perforated membrane having a plurality of preselected perforated portions each holding and dispensing a reproducible unit dose of less than 50 mg of the said inhalable pwder, said membrane being mounted for movement between said conduit and said storage chamber so that one of said preselected portions is positioned across said conduit whereby the active compound held in the perforation thereof can be dispensed into the conduit and another of said preselected portions thereof is disposed within said storage chamber, dose loading means for introducing said inhalable powder in the storage chamber into the perforation of the preselected portion of said membrane disposed within the

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storage chamber, and maneuvering means for displacing the perforated membrane through a plurality of positions whereby successive preselected portions of the perforated membrane holding the inhalable powder are positioned across said conduit for dispensing the inhalable powder.

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In another embodiment, the invention relates to a method for treatment of airway diseases, particularly COPD (chronic obstructive pulmonary disease) and asthma, charcterized in that an inhalable powder containing tiotropium, preferably in an \_amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with urraverage particle size of between 10 to 500 µm, is administered via inhalation by the inhaler according to US 4,524,769, comprising a nozzle, a conduit connected to the nozzle, a storage chamber adjacent said conduit for storing said inhalable powder to be dispensed by said inhalator. a perforated membrane having a plurality of preselected perforated portions each holding and dispensing a reproducible unit dose of less than 50 mg of the said inhalable powder, said membrane being mounted for movement between said conduit and said storage chamber so that one of said preselected portions is positioned across said conduit whereby the active compound held in the perforation thereof can be dispensed into the conduit and another of said preselected portions thereof is disposed within said storage chamber, dose loading means for introducing said inhalable powder in the storage chamber into the perforation of the preselected portion of said membrane disposed within the storage chamber, and maneuvering means for displacing the perforated membrane through a plurality of positions whereby successive preselected portions of the perforated membrane holding the inhalable powder are positioned across said conduit for dispensing the inhalable powder.

In another preferred embodiment the invention relates to the use of the inhaler according to US 4,524,769 comprising

a nozzle, a conduit connected to the nozzle, a storage chamber adjacent said conduit for storing said inhalable powder to be dispensed by said inhalator, a perforated membrane having a plurality of preselected perforated portions each holding and dispensing a reproducible unit dose of less than 50 mg of the said inhalable pwder, said membrane being mounted for movement between said conduit and said storage chamber so that one of said preselected portions is positioned across said conduit whereby the active compound held in the perforation thereof can be dispensed into the conduit and another of said preselected portions thereof is disposed within said storage chamber, dose loading means for introducing said inhalable powder in the storage chamber into the perforation of the preselected

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portion of said membrane disposed within the storage chamber, and maneuvering means for displacing the perforated membrane through a plurality of positions whereby successive preselected portions of the perforated membrane holding the inhalable powder are positioned across said conduit for dispensing the inhalable powder,

for the administration of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m.

In yet another preferred embodiment the invention relates to an inhalation kit consisting of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 μm, and the inhaler according to US 4,524,769, comprising

a nozzle, a conduit connected to the nozzle, a storage chamber adjacent said conduit for storing said inhalable powder to be dispensed by said inhalator, a perforated membrane having a plurality of preselected perforated portions each holding and dispensing a reproducible unit dose of less than 50 mg of the said inhalable pwder, said membrane being mounted for movement between said conduit and said storage chamber so that one of said preselected portions is positioned across said conduit whereby the active compound held in the perforation thereof can be dispensed into the conduit and another of said preselected portions thereof is disposed within said storage chamber, dose loading means for introducing said inhalable powder in the storage chamber into the perforation of the preselected portion of said membrane disposed within the storage chamber, and maneuvering means for displacing the perforated membrane through a plurality of positions whereby successive preselected portions of the perforated membrane holding the inhalable powder are positioned across said conduit for dispensing the inhalable powder.

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In another preferred embodiment according to the invention the inhaler according to US 5,590,645 is applied. The disclosure of US 5,590,645 is incorporated herein by reference in its entirety.

Accordingly, in a preferred embodiment the invention relates to a method for the administration of an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 μm, by means of the inhaler according to 5,590,645, comprising

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a medicament pack having a plurality of containers for containing medicament in powder form wherein the containers are spaced along the length of and defined between two peelable sheets secured to each other, an opening station for receiving a container of said medicament pack being, means positioned to engage peelable sheets of a container which has been received in said opening station for peeling apart the peelable sheets, to open such a container, an outlet, positioned to be in communication with an opened container, through which a user can inhale medicament in powder form from such an opened container, and indexing means for indexing in communication with said outlet containers of a medicament pack in use with said inhalation device.

In another embodiment, the invention relates to a method for treatment of airway diseases, particularly COPD (chronic obstructive pulmonary disease) and asthma. charcterized in that an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with 15 an average particle size of between 10 to 500 µm, is administered via inhalation by the inhaler according to US 5,590,645, comprising a medicament pack having a plurality of containers for containing medicament in powder form wherein the containers are spaced along the length of and defined between two peelable sheets secured to each other, an opening station for receiving 20 a container of said medicament pack being, means positioned to engage peelable sheets of a container which has been received in said opening station for peeling apart the peelable sheets, to open such a container, an outlet, positioned to be in communication with an opened container, through which a user can inhale medicament in powder form from such an opened container, and indexing means for indexing in communication with said outlet containers of a medicament pack in use with said inhalation device.

In another preferred embodiment the invention relates to the use of the inhaler according to US 5,590,645, comprising a medicament pack having a plurality of containers for containing medicament in powder form wherein the containers are spaced along the length of and defined between two peelable sheets secured to each other, an opening station for receiving a container of said medicament pack being, means positioned to engage peelable sheets of a container which has been received in said opening station for peeling apart the peelable sheets, to open such a container, an outlet, positioned to be in communication with an opened container, through which a user can inhale medicament in powder form from such an opened container, and

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indexing means for indexing in communication with said outlet containers of a medicament pack in use with said inhalation device,

for the administration of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm.

In yet another preferred embodiment the invention relates to an inhalation kit consisting of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an . 10 average particle size of between 10 to 500 µm, and the inhaler according to US 5.590.645, comprising a medicament pack having a plurality of containers for containing medicament in powder form wherein the containers are spaced along the length of and defined between two peelable sheets secured to each other, an opening station for receiving a container of said medicament pack being, means positioned to engage peelable 15 sheets of a container which has been received in said opening station for peeling apart the peelable sheets, to open such a container, an outlet, positioned to be in communication with an opened container, through which a user can inhale medicament in powder form from such an opened container, and indexing means for indexing in communication with said outlet containers of a 20 medicament pack in use with said inhalation device.

In another preferred embodiment according to the invention the inhaler according to US 4,627,432 is applied. The disclosure of US 4,627,432 is incorporated herein by reference in its entirety.

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Accordingly, in a preferred embodiment the invention relates to a method for the administration of an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m, by means of the inhaler according to US 4,627,432, being characterised by a housing with a chamber therein, an air inlet into the chamber.

a circular disc having an axis substantially coaxial to the chamber axis and rotatable inside the chamber and provided with a plurality of apertures therethrough arranged in a circle, said apertures being sized and positioned so that each aperture is adapted to be aligned with a different container, the said disc being arranged so that the carrier can be placed in contact with one face of the disc with one of the containers located in each one of the apertures, an outlet through which a patient may inhale leading out of the chamber, an opening in said housing alignable with

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respective ones of the apertures in the disc as the disc is rotated, a plunger operatively connected to said housing and having a penetrating member, said penetrating member being displaceable to pass through said opening and the corresponding aperture in the disc registered with it thereby to penetrate and open a container located in the aperture so that the medicament will be released from the container and entrained in the air flow produced by a patient inhaling through the outlet, and means between said disc and said housing for rotatably indexing the disc to register each of the apertures in turn with the housing opening.

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In another embodiment, the invention relates to a method for treatment of airway 10 diseases, particularly COPD (chronic obstructive pulmonary disease) and asthma. charcterized in that an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm, is administered via inhalation by 15 the inhaler according to US 4,627,432, being characterised by a housing with a chamber therein, an air inlet into the chamber, a circular disc having an axis substantially coaxial to the chamber axis and rotatable inside the chamber and provided with a plurality of apertures therethrough arranged in a circle, said apertures being sized and positioned so that each aperture is adapted to be aligned with a different container, the said disc being arranged so that 20 the carrier can be placed in contact with one face of the disc with one of the containers located in each one of the apertures, an outlet through which a patient may inhale leading out of the chamber, an opening in said housing alignable with respective ones of the apertures in the disc as the disc is rotated, a plunger operatively connected to said housing and having a penetrating member, said 25 penetrating member being displaceable to pass through said opening and the corresponding aperture in the disc registered with it thereby to penetrate and open a container located in the aperture so that the medicament will be released from the container and entrained in the air flow produced by a patient inhaling through the outlet, and means between said disc and said housing for rotatably indexing the disc 30 to register each of the apertures in turn with the housing opening.

In another preferred embodiment the invention relates to the use of the inhaler according to US 4,627,432 being characterised by a housing with a chamber therein, an air inlet into the chamber,

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a circular disc having an axis substantially coaxial to the chamber axis and rotatable inside the chamber and provided with a plurality of apertures therethrough arranged in a circle, said apertures being sized and positioned so that each aperture is adapted to be aligned with a different container, the said disc being arranged so that

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the carrier can be placed in contact with one face of the disc with one of the containers located in each one of the apertures, an outlet through which a patient may inhale leading out of the chamber, an opening in said housing alignable with respective ones of the apertures in the disc as the disc is rotated, a plunger operatively connected to said housing and having a penetrating member, said penetrating member being displaceable to pass through said opening and the corresponding aperture in the disc registered with it thereby to penetrate and open a container located in the aperture so that the medicament will be released from the container and entrained in the air flow produced by a patient inhaling through the outlet, and means between said disc and said housing for rotatably indexing the disc to register each of the apertures in turn with the housing opening, for the administration of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 µm.

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In yet another preferred embodiment the invention relates to an inhalation kit consisting of an inhalable powdered containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m, and the inhaler according to US 4,627,432, being characterised by a housing with a chamber therein, an air inlet into the chamber,

a circular disc having an axis substantially coaxial to the chamber axis and rotatable inside the chamber and provided with a plurality of apertures therethrough arranged in a circle, said apertures being sized and positioned so that each aperture is adapted to be aligned with a different container, the said disc being arranged so that the carrier can be placed in contact with one face of the disc with one of the containers located in each one of the apertures, an outlet through which a patient may inhale leading out of the chamber, an opening in said housing alignable with respective ones of the apertures in the disc as the disc is rotated, a plunger operatively connected to said housing and having a penetrating member, said penetrating member being displaceable to pass through said opening and the corresponding aperture in the disc registered with it thereby to penetrate and open a container located in the aperture so that the medicament will be released from the container and entrained in the air flow produced by a patient inhaling through the outlet, and means between said disc and said housing for rotatably indexing the disc to register each of the apertures in turn with the housing opening.

The following Examples serve to illustrate the present invention further without restricting its scope to the embodiments provided hereinafter by way of example.

# Starting materials

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As a starting material for the synthesis of crystalline tiotropiumbromide monohydrate tiotropiumbromide obtained according to the disclosure of European patent application EP 418 716 A1 is be used.

# Preparation of crystalline tiotropium bromide monohydrate:

15.0 kg of tiotropium bromide as obtained according to the methods disclosed in EP 418 716 A1 are added to 25.7 kg of water in a suitable reaction vessel. The mixture is heated to 80-90°C and stirred at constant temperature until clear solution is formed. Activated charcoal (0.8 kg), moistened with water, is suspended in 4.4 kg of water, this mixture is added to the solution containing the tiotropium bromide and rinsed with 4.3 kg of water. The mixture thus obtained is stirred for at least 15 min at 80-90°C and then filtered through a heated filter into an apparatus which has been preheated to an outer temperature of 70°C. The filter is rinsed with 8.6 kg of water. The contents of the apparatus are cooled at 3-5°C every 20 minutes to a temperature of 20-25°C. The apparatus is further cooled to 10-15°C using cold water and crystallisation is completed by stirring for at least one hour. The crystals are isolated using a suction drier, the crystal slurry isolated is washed with 9 litres of cold water (10-15°C) and cold acetone (10-15°C). The crystals obtained are dried in a nitrogen current at 25°C over 2 hours.

Yield: 13.4 kg of tiotropium bromide monohydrate (86 % of theory)

The crystalline tiotropium bromide monohydrate thus obtained is micronised by known methods, to bring the active substance into the average particle size which meets the specifications according to the invention.

The DSC diagram of crystalline tiotropium bromide monohydrate shows two characteristic signals. The first, relatively broad, endothermic signal between 50-120°C can be attributed to the dehydration of the tiotropium bromide monohydrate to produce the anhydrous form. The second, relatively sharp endothermic peak at 230  $\pm$  5°C can be put down to the melting of the substance. These data were obtained using a Mettler DSC 821 and evaluated with the Mettler STAR software package. These data, like the other values given in the above Table, were obtained at a heating rate of 10 K/min.

The crystalline tiotropium bromide monohydrate thus obtained was characterised by IR spectroscopy. The data was obtained using a Nicolet FTIR spectrometer and

evaluated with the Nicolet OMNIC software package, version 3.1. The measurement was carried out with 2.5µmol of tiotropium bromide monohydrate in 300 mg of KBr. Table 1 shows some of the essential bands of the IR spectrum.

5	Table	1:	<b>Attribution</b>	of s	pecific	bands
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_	Wave number (cm <sup>-1</sup> )	Attribution	Type of oscillation
	3570, 410	О-Н	elongated
			oscillation
٠.,	3105	Aryl C-H	elongated
•		·	oscillation
	1730	C=O	elongated
			oscillation
	1260	Epoxide C-O	elongated
			oscillation
	1035	Ester C-OC	elongated
			oscillation
	720	Thiophene	cyclic oscillation

The crystalline tiotropium bromide monohydrate was characterised by X-ray structural analysis. The measurements of X-ray diffraction intensity were carried out on an AFC7R- 4-circuit diffractometer (Rigaku) using monochromatic copper  $K_{\alpha}$  radiation. The structural solution and refinement of the crystal structure were obtained by direct methods (SHELXS86 Program) and FMLQ-refinement (TeXsan Program). The X-ray structural analysis carried out showed that crystalline tiotropium bromide hydrate has a simple monoclinic cell with the following dimensions: a = 18.0774 Å, b = 11.9711 Å, c = 9.9321 Å,  $\beta = 102.691^{\circ}$ ,  $V = 2096.96 \text{ Å}^3$ .

### **Apparatus**

The following machines and equipment, for example, may be used to prepare the inhalable powders according to the invention:

Mixing container or powder mixer: Gyrowheel mixer 200 L; type: DFW80N-4; made by: Messrs Engelsmann, D-67059 Ludwigshafen.

25 Granulating sieve: Quadro Comil; type: 197-S; made by: Messrs Joisten & Kettenbaum, D-51429 Bergisch-Gladbach.

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The following examples provide for inhalable powder mixtures applicable according to the invention.

# Example 1:

5.2 kg of glucose monohydrate for inhalation (average particle size 25µm) are used as the excipient. 22.5 g crystalline tiotropiumbromide monohydrate (micronised; average particle size 1 - 3.5 µm) are used as the active ingredient.

The aforementioned components are sieved in in alternate layers of lactose monohydrate in batches of about 200 g and crystalline tiotropiumb ലേവര് 👉 monohydrate in batches of about 1g. The ingredients sieved in are then mixed together (mixing at 900 rpm).

According to the invention preferably 5.2225 mg of the aforementioned powder are delivered per dose. 15

# Example 2:

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5.4775 kg of lactose monohydrate for inhalation (average particle size 25µm) are used as the excipient. 22.5 g crystalline tiotropiumbromide monohydrate (micronised; average particle size 1 - 3.5 µm) are used as the active ingredient.

The aforementioned components are sieved in in alternate layers of lactose monohydrate in batches of about 200 g and crystalline tiotropiumbromide monohydrate in batches of about 1g. The ingredients sieved in are then mixed 25 together (mixing at 900 rpm).

According to the invention preferably 5.5 mg of the aforementioned powder are delivered per dose.

#### 30 Example 3:

### 1.1: Excipient mixture:

5.203 kg of lactose monohydrate for inhalation (average particle size 25 µm) are used as the coarser excipient component. 0,27 kg of lactose monohydrate (5µm) are used as the finer excipient component. In the resulting 5,473 kg of excipient mixture the proportion of the finer excipient component is 5%.

The aforementioned components are sieved in in alternate layers of lactose monohydrate (25 µm) in batches of about 200 g and lactose monohydrate (5 µm) in

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batches of about 10g. The ingredients sieved in are then mixed together (mixing at 900 rpm).

# 1.2: Final mixture:

To prepare the final mixture, 5,473 kg of the excipient mixture (1.1) and 22.5 g crystalline tiotropiumbromide monohydrate (micronised; average particle size 1 - 3.5 μm) are used. The content of active substance in the resulting powder is 0.4%.

The aforementioned components are sieved in in alternate layers of excipient mixture (1.1) in batches of about 200 g and crystalline tiotropiembromide monohydrate in batches of about 1g. The ingredients sieved in are then mixed together (mixing at 900 rpm).

According to the invention preferably about 5.5 mg of the aforementioned powder are delivered per dose.

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### **Patent Claims**

- Use of an inhaler for the administration of a tiotropium containing inhalable powder via inhalation, characterised in that the inhalable powder contains tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500 μm, and further characterized in that the said inhaler displays a flow resistance of about 0.01 0.1 √kPa min/l.
- 10 2) Use according to claim 1, characterised in that the inhaler is characterized by a flow resistance of about  $0.02 0.06 \sqrt{kPa}$  min/l.
- 3) Use according to claim 1 or 2, characterised in that the inhaler comprises a housing, containing two windows, a deck in which there are air inlet ports and which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring, a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in the central region around the capsule chamber and underneath the screen housing and screen.
  - 4) Use according to one of claims 1, 2 or 3, characterised in that tiotropium is used in form of its chloride, bromide, iodide, methanesulphonate, paratoluenesulphonate or methyl sulphate, preferably in form of its bromide.
  - 5) Use according to claim 4, characterised in that tiotropium is used in form of its crystalline tiotropium bromide monohydrate.
- Inhalation kit consisting of an inhaler displaying a flow resistance of about 0.01  $-0.1 \sqrt{kPa}$  min/l and an inhalable powder containing tiotropium, preferably in an amount of 0.001 to 5 %, in admixture with a physiologically acceptable excipient with an average particle size of between 10 to 500  $\mu$ m.
- Inhalation kit according to claim 6, characterised in that the inhaler is characterized by a flow resistance of about  $0.02 0.06 \sqrt{kPa}$  min/l.

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8) Inhalation kit according to claim 6 or 7, characterised in that the inhaler comprises a housing, containing two windows, a deck in which there are air inlet ports and which is provided with a screen secured via a screen housing, an inhalation chamber connected to the deck on which there is a push button provided with two sharpened pins and movable counter to a spring, a mouthpiece which is connected to the housing, the deck and a cover via a spindle to enable it to be flipped open or shut, and three holes with diameters below 1 mm in the central region around the capsule chamber and underneath the screen housing and screen.

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9) Inhalation kit according to one of claims 6, 7 or 8, characterised in that tiotropium is present in form of its chloride, bromide, iodide, methanesulphonate, para-toluenesulphonate or methyl sulphate, preferably in form of its bromide.

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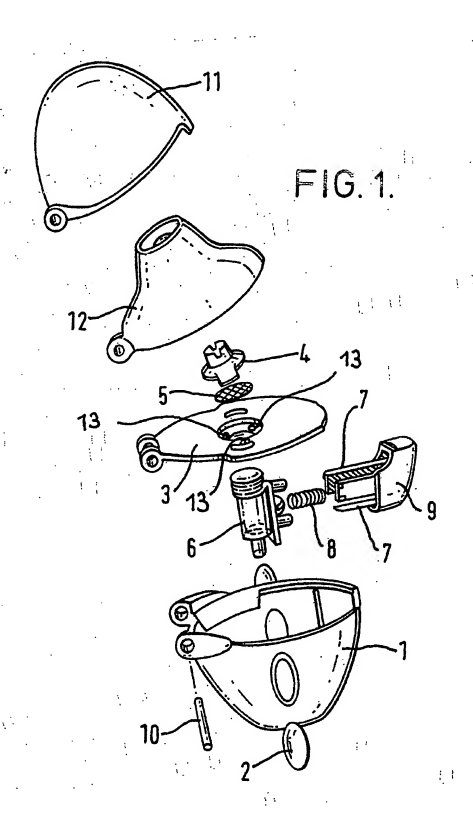
10) Inhalation kit according to claim 9, characterised in that tiotropium is present in form of its crystalline tiotropium bromide monohydrate.

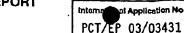
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A. CLASSIFICATION OF SUBJECT MATTER IPC 7 A61K9/00 A61k A61K31/46 A61M15/00 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC 7-A61K-A61MDocumentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicel, search terms used) EPO-Internal, WPI Data, PAJ, EMBASE, BIOSIS C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Category ° 1-10 χ WO OO 28979 A (SKYEPHARMA AG ;MUELLER WALZ RUDI (DE); KELLER MANFRED (DE)) 25 May 2000 (2000-05-25) page 18, line 26 -page 18, line 31; example 6 1 - 10X EP 1 158 970 A (NOVARTIS ERFIND VERWALT GMBH : NOVARTIS AG (CH)) 5 December 2001 (2001-12-05) example 3 US 5 590 645 A (DAVIES MICHAEL BIRSHA ET 1 - 10AL) 7 January 1997 (1997-01-07) figures 1-34 US 4 627 432 A (NEWELL ROBERT E ET AL) 1-10 Y 9 December 1986 (1986-12-09) figures 1-4 Further documents are listed in the continuation of box C. Patent family members are listed in annex. l X I Special categories of cited documente : 'T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the \*A\* document defining the general state of the art which is not considered to be of particular relevance invention \*E\* earlier document but published on or after the international filing date "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to hyotre an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled "O" document referring to an oral disclosure, use, exhibition or \*P\* document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 25/07/2003 14 July 2003 Authorized officer Name and mailing address of the ISA European Palent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nt, ESTANOL, I

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International Application No PCT/EP 03/03431

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Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	mational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.:  because they relate to subject matter not required to be searched by this Authority, namely:
	Rule 39.1(1v) PCT — Method for treatment of the human or animal body by therapy
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
з. 🗌	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This inte	mational Searching Authority found multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3.	As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest  The additional search fees were accompanied by the applicant's protest.  No protest accompanied the payment of additional search fees.

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